GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: June 15, 2005, 10:19:12; Search time 178.268 Seconds

(without alignments)

846.121 Million cell updates/sec

Title: US-10-053-975A-1

Perfect score: 2047

Sequence: 1 MAVSESQLKKMVSKYKYRDL.....FQLRALMQKARKTAGLSDLY 390

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2105692 segs, 386760381 residues

Total number of hits satisfying chosen parameters: 2105692

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : A_Geneseq_16Dec04:*

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4: geneseqp2001s:*

5: geneseqp2002s:*

6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		8			·	
Result		Query				
No.	Score	Match	Length	DB	ID	Description
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1				_		
2	2047	100.0	390	5	ABP52191	Abp52191 Human tum
3	2047	100.0	390	6	ABG73787	Abg73787 Human tum
4	2047	100.0	390	7	ADC35176	Adc35176 Novel hum
5	2047	100.0	390	8	ADF91425	Adf91425 TSG101 #S
6	2047	100.0	390	8	ADL82957	Adl82957 Human PRO
7	2047	100.0	390	8	ADN12256	Adn12256 Human TSG
8	2047	100.0	390	8	ADO43235	Ado43235 Human TSG
9	2047	100.0	390	8	ADO43233	Ado43233 Human TSG

10	2047	100.0	390	8	ADP82614		Adp82614 Human tum
11		100.0	390	8	ADP56114		Adp56114 Human PRO
12	2042	99.8	390	6	AAE34885		Aae34885 Human Tsg
13	2002	97.8	380	2	AAW19111		Aaw19111 Human tum
14	1945.5	95.0	391	٠6	AAE34884		Aae34884 Mouse Tsg
15		95.0	391	7	ADB85228		Adb85228 Mouse tum
16		92.8	381	2	AAW19110		Aaw19110 Mouse tum
17		92.8	381	2	AAW93425		Aaw93425 Mouse tsg
18	1900.5	92.8	381	6	ABG73788		Abg73788 Murine ts
19	1900.5	92.8	381	7	ADC35178		Adc35178 Human tsg
20	1404	68.6	307	5	ABP41729		Abp41729 Human ova
21	1162	56.8	237	4	AAE09328		Aae09328 Human int
22	809.5	39.5	331	4	ABB64607		Abb64607 Drosophil
23	776	37.9	145	8	ADF91424		Adf91424 TSG101UEV
. 24	479	23.4	90	3	AAG01689		Aag01689 Human sec
25	455	22.2	398	3	AAG06370		Aag06370 Arabidops
26	455	22.2	398	8	ADN73283		Adn73283 Thale cre
27	455	22.2	412	3	AAG06369		Aag06369 Arabidops
28	435	21.3	379	5	ABP43551		Abp43551 L-lactate
29		21.3	379	8	ADH13697		Adh13697 Human ENZ
30		20.7	340	3	AAG06371		Aag06371 Arabidops
31	390	19.1	87	8	AB055273		Abo55273 Human gen
32	380	18.6	322	3	AAG13898		Aag13898 Arabidops
33	371	18.1	341	4	AAB93473		Aab93473 Human pro
34		18.1	433	7	ADM29340		Adm29340 Human nov
35	346	16.9	146	3	AAB53717		Aab53717 Human col
36	339	16.6	452	4	AAU23116		Aau23116 Novel hum
37	328	16.0	73	8	ABO55495		Abo55495 Human gen
38	267.5	13.1	288	3	AAG13899		Aag13899 Arabidops
39	161.5	7.9	168	3	AAG13900		Aag13900 Arabidops
40	161	7.9	485	3	AAB57086		Aab57086 Human pro
41	160	7.8	466	6	ABO07215		Abo07215 Human p53
42	160	7.8	466	7	ADD47589		Add47589 Human Pro
43	160	7.8	466	7	ADI15879		Adi15879 Human PP
44	160	7.8	466	7	ADJ68571		Adj68571 Human hea
45	153	7.5	218	4	ABG18833		Abg18833 Novel hum
46	151	7.4	466	2	AAR07084		Aar07084 Recombina
47	148.5	7.3	148	3	AAY86515		Aay86515 Human gen
48	148.5	7.3	148	5	ABB97431		Abb97431 Novel hum
49	148.5	7.3	148	6	AB053679		Abo53679 Novel hum
. 50		7.3	148	8	ABM81395		Abm81395 Tumour-as
51		7.3	176	4	AAG75132		Aag75132 Human col
52		7.3	397	4	AAM39852		Aam39852 Human pol
- 53	148.5	7.3	397	4	AAG81381		Aag81381 Human AFP
54	148.5	7.3	397	6	ADA55415		Ada55415 Human pro
55	148.5	7.3	442	4	AAM41638		Aam41638 Human pol
56		7.1	368	8	ADM97711	·	Adm97711 Human MNA
57		7.1	536	8	ADM97710		Adm97710 Human MNA
58		7.1	1130	8	ADM97695		Adm97695 Human MNA
59		7.1	777	5	ABP41839		Abp41839 Human ova
60		7.1	1130	7	ADC99065		Adc99065 Human KPP
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62		6.9	505	2	AAY07117		Aay07117 Lung canc
63		6.9	505	6	AB007216		Abo07216 Human p53
64		6.9	505	7	ADJ68733		Adj68733 Human hea
65		6.9	505	8	ABM81156		Abm81156 Tumour-as
66	142	6.9	505	8	ADP23965		Adp23965 PRO polyp

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67	142	6.9	905	2	AAW31186	Aaw31186 Human p16
68	142	6.9	983	8	ADM87342	Adm87342 Human pro
69	142	6.9	1021	7	ADJ68507	Adj68507 Human hea
70	142	6.9	1135	2	AAW31185	Aaw31185 Human p16
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72	142	6.9	1251	8	ADM87788	Adm87788 Human EST
73	141	6.9	244	3	AAG40015	Aag40015 Arabidops
74	141	6.9	245	3	AAG38059	Aag38059 Arabidops
75	141	6.9	245	3	AAG05889	Aag05889 Arabidops
76	141	6.9	247	3	AAG38058	Aag38058 Arabidops
77	141	6.9	247	3	AAG05888	Aag05888 Arabidops
78	141	6.9	263	3	AAG38057	Aag38057 Arabidops
79	141	6.9	274	3	AAG05887	Aag05887 Arabidops
80	141	6.9	463	7	ADD47587	Add47587 Rát Prote
81	140.5	6.9	1277	4	ABB66688	Abb66688 Drosophil
82	139.5	6.8	258	3	AAB44500	Aab44500 Plant vir
83	138.5	6.8	827	5	ABP35602	Abp35602 Fungal ZB
84	135.5	6.6	177	4	ABG09342	Abg09342 Novel hum
85	135	6.6	121	3	AAG21511	Aag21511 Arabidops
86	134.5	6.6	172	4	ABG18832	Abg18832 Novel hum
87	133	6.5	144	4	AAM42061	Aam42061 Human pol
88	132	6.4	369	7	ADD14178	Add14178 Human src
89	132	6.4	371	2	AAW64535	Aaw64535 Human leu
90	132	6.4	371	5	ABP61444	Abp61444 Human NF-
91	132	6.4	371	6	ABR41017	Abr41017 Human MAP
92	132	6.4	371	8	ADR09413	Adr09413 Human pro
93	131	6.4	101	3	AAG21512	Aag21512 Arabidops
94	131	6.4	518	8	ADN20587	Adn20587 Bacterial
95	131	6.4	539	3	AAB10224	Aab10224 Chicken n
96	130.5	6.4	237	6	ABR58292	Abr58292 BCU0371B
97	130.5	6.4	338	· 7	ADB75310	Adb75310 Prostate
98	130.5	6.4	338	8	ADK70560	Adk70560 Respirato
99	130.5	6.4	338	8	ADQ17947	Adq17947 Human sof
100	130.5	6.4	338	8	ADP23414	Adp23414 PRO polyp

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RESULT 1
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ID
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AC
XX
     11-JUN-1999 (first entry)
DT
XX
     Human NHTS protein.
DE
XX
KW
     Human tumour suppressor; NHTS; treatment; diagnosis; lymphoma; cancer;
     brain; breast; colon; heart; kidney; ovary; paraganglia; pancreas;
KW
     prostate; skin; stomach; thyroid; autoimmune disease; asthma; diabetes;
KW
     biliary cirrhosis; Crohn's disease; rheumatoid arthritis.
KW
XX
     Homo sapiens.
os
XX
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PN
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XX
PD
    06-APR-1999.
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    23-JAN-1997;
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XX
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                 97US-00786999.
PR
XX
    (INCY-) INCYTE PHARM.
PA
XX
PΙ
    Brie SL, Goli SK;
XX
    WPI; 1999-253932/21.
DR
    N-PSDB; AAX23168.
DR
XX
    Novel human tumor suppressor - useful for the diagnosis or treatment of
PT
PT
    lymphoma, cancer, and autoimmune disease.
XX
    Claim 1; Fig 1A-B; 25pp; English.
PS
XX
    This invention describes the isolation of a novel human tumour suppressor
CC
    (NHTS). The products of the invention may be used for the diagnosis or
CC
    treatment of conditions and diseases which are associated with expression
CC
    of NHTS e.g. lymphoma and cancers of the brain, breast, colon, heart,
CC
    kidney, ovary, paraganglia, pancreas, prostate, skin, stomach and thyroid
CC
    and autoimmune disease e.g. asthma, biliary cirrhosis, Crohn's disease,
CC
CC
    diabetes, and rheumatoid arthritis
XX
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                                              Length 390;
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                      100.0%; Pred. No. 7e-149;
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                                                       0;
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                                                          Gaps
                                                                 0;
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Db
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Qν
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Db
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us-10-053-975a-1.rai

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

June 15, 2005, 10:33:17; Search time 46.8487 Seconds Run on:

(without alignments)

621.429 Million cell updates/sec

US-10-053-975A-1 Title:

Perfect score: 2047

1 MAVSESQLKKMVSKYKYRDL.....FQLRALMQKARKTAGLSDLY 390 Sequence:

Scoring table: BLOSUM62

Gapop 10.0, Gapext 0.5

513545 segs, 74649064 residues Searched:

Total number of hits satisfying chosen parameters: 513545

Minimum DB seq length: 0

Maximum DB sed length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 100 summaries

Issued_Patents_AA:* Database:

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			%				• •
R	esult No.	Score	Query	Length	DB	ID	Description
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	4 5 6 7	2047 2002 2002 2002	100.0 97.8 97.8 97.8	403 380 380 380	4 1 2	US-09-949-016-11251 US-08-585-758A-4 US-08-977-818-4 US-08-670-274B-4	Sequence 11251, A Sequence 4, Appli Sequence 4, Appli Sequence 4, Appli
	8 9 10 11	2002 2002 1945.5 1900.5	97.8 97.8 95.0 92.8	380 380 391 381	3 4 4 1	US-09-146-187-4 US-09-804-690-4 US-09-886-319A-1 US-08-585-758A-2	Sequence 4, Appli Sequence 4, Appli Sequence 1, Appli Sequence 2, Appli
	12 13 14 15 16 17	1900.5 1900.5 1900.5 1900.5 1900.5	92.8 92.8 92.8 92.8 92.8 92.8	381 381 381 381 381	1 2 2 3 4 4	US-08-977-818-2 US-08-670-2748-2 US-08-786-999-3 US-09-146-187-2 US-09-216-387-3 US-09-804-690-2	Sequence 2, Appli Sequence 2, Appli Sequence 3, Appli Sequence 2, Appli Sequence 3, Appli Sequence 2, Appli
		1500.5	32.0	301	•	05 05 00. 050 2	requested 21 Appil

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81	120	5.9	777	4	US-09-949-016-8707	Sequence 8707, Ap
82	119.5	5.8	503	3	us-08-526-136-2	Sequence 2, Appli
83	119.5	5.8	505	3	us-08-526-136-4	Sequence 4, Appli
84	118.5	5.8	397	4	US-09-248-796A-18882	Sequence 18882, A
85	118.5	5.8	1255	2	us-09-080-897-4	Sequence 4, Appli
86	118.5	5.8	1255	3	US-08-899-595-1	Sequence 1, Appli
87	118.5	5.8	1255 .	3	US-09-323-735-4	Sequence 4, Appli
88	118	5.8	214	1	US-08-217-327-4	Sequence 4, Appli
89	118	5.8	581	4	us-09-949-016-9978	Sequence 9978, Ap
90	117 ·	5.7	416	4	US-09-690-454-136	Sequence 136, App
91	117	5.7	779	4	US-10-164-595-56	Sequence 56, Appl
92	117	5.7	843	4	US-10-164-595-54	Sequence 54, Appl
93	116.5	5.7	338	1	US-08-218-686-2	Sequence 2, Appli
94	116.5	5.7	338	3	us-08-460-242-2	Sequence 2, Appli
95	116	5.7	1461	4	us-09-585-887-9	Sequence 9, Appli
96	116	5.7	1461	4	US-09-289-578-9	Sequence 9, Appli
97	116	5.7	1464	4	US-09-331-347C-21	Sequence 21, Appl
98	115.5	5.6	357	1	us-07-609-716-66	Sequence 66, Appl
99	115.5	5.6	357	1	us-08-642-255-33	Sequence 33, Appl
100	115.5	5.6	357	3	US-08-475-411A-66	Sequence 66, Appl

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RESULT 1
us-08-786-999-1
  Sequence 1, Application US/08786999 Patent No. 5892016
   GENERAL INFORMATION:
     APPLICANT:
                  La Brie, Sam
     APPLICANT:
                   Goli, Surya K.
     TITLE OF INVENTION: NOVEL HUMAN TUMOR
     TITLE OF INVENTION:
                             SUPPRESSOR
     NUMBER OF SEQUENCES:
     CORRESPONDENCE ADDRESS:
        ADDRESSEE: Incyte Pharmaceuticals, Inc. STREET: 3174 Porter Drive
        CITY: Palo Alto
STATE: CA
COUNTRY: USA
        ZIP: 94304
     COMPUTER READABLE FORM:
        MEDIUM TYPE: Diskette
        COMPUTER: IBM Compatible
        OPERATING SYSTEM: DOS
                    FastSEQ for Windows Version 2.0
        SOFTWARE:
     CURRENT APPLICATION DATA:
        APPLICATION NUMBER: US/08/786,999
        FILING DATE: Filed Herewith
        CLASSIFICATION: 530
      PRIOR APPLICATION DATA:
        APPLICATION NUMBER:
        FILING DATE:
     ATTORNEY/AGENT INFORMATION:
        NAME: Billings, Lucy J.
REGISTRATION NUMBER: 36,749
REFERENCE/DOCKET NUMBER: PF-0199 US
     TELECOMMUNICATION INFORMATION:
        TELEPHONE: 415-855-0555
        TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO:
      SEQUENCE CHARACTERISTICS:
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us-10-053-975a-1.rai

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LENGTH: 390 amino acids
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     STRANDEDNESS: single
     TOPOLOGY:
              linear
    IMMEDIATE SOURCE:
     LIBRARY: Colnnot01
CLONE: 609476
ÚS-08-786-999-1
                     100.0%; Score 2047; DB 2; 100.0%; Pred. No. 7.3e-159;
                                             Length 390:
 Query Match
                     100.0%;
 Best Local Similarity
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                                                               0:
 Matches 390: Conservative
                                             Indels
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           Db
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Qy
           Db
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us-10-053-975a-1.rapb

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OM protein - protein search, using sw model

Run on:

June 15, 2005, 10:46:54; Search time 160.016 Seconds

(without alignments)

934.287 Million cell updates/sec

Title:

US-10-053-975A-1

Perfect score:

2047

Sequence:

1 MAVSESQLKKMVSKYKYRDL......FQLRALMQKARKTAGLSDLY 390

Scoring table:

BLOSUM62

Gapop 10.0, Gapext 0.5

Searched:

1710399 seqs, 383334425 residues

Total number of hits satisfying chosen parameters:

1710399

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database :

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19: /cgn2_6/ptodata/1/pubpaa/US11_NEW_PUB.pep:* 20:

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description	
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; Publication No. US20020086019A1
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; APPLICANT: Wolf, Eckard
; APPLICANT: Werner, Sabine
; APPLICANT: Werner, Sabine
; APPLICANT: Regenbogen, Johannes
; APPLICANT: Regenbogen, Johannes
; APPLICANT: Goppelt, Andreas
; TITLE OF INVENTION: Use of Polypeptides or Nucleic Acids for
; TITLE OF INVENTION: the Diagnosis or Treatment of Skin Disorders and Wound
; TITLE OF INVENTION: Healing and for the Identification of Pharmacologically
; TITLE OF INVENTION: Active Substances
; FILE REFERENCE: 50125/014002
; CURRENT APPLICATION NUMBER: US/09/886,319A
; CURRENT FILING DATE: 2001-06-20
; PRIOR APPLICATION NUMBER: US 60/222,081
; PRIOR FILING DATE: 2000-08-01
; PRIOR APPLICATION NUMBER: DE 10030149.5
; PRIOR FILING DATE: 2000-06-20
; NUMBER OF SEQ ID NOS: 84
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Db

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: June 15, 2005, 10:32:03; Search time 36.5055 Seconds

(without alignments)

1027.916 Million cell updates/sec

Title: US-10-053-975A-1

Perfect score: 2047

Sequence: 1 MAVSESQLKKMVSKYKYRDL......FQLRALMQKARKTAGLSDLY 390

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 segs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : PIR_79:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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transforming prote	3	TVHUJB	1	347	5.5	113.5	93
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Science 282, 2012-2018, 1998
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investigating biology.
A; Reference number: A75000; MUID: 99069613; PMID: 9851916
A; Note: see websites genome.wustl.edu/gsc/C elegans/ and
www sanger.ac.uk/Projects/C_elegans/ for a list of authors
A; Note: published errata appeared in Science 283, 35, 1999; Science 283, 2103,
1999; and Science 285, 1493, 1999
A; Accession: H88650
A; Status: preliminary
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A;Cross-references: UNIPROT:076258; GB:chr_IV; PIDN:AAC25822.1; PID:g3294495;
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C; Genetics:
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A; Gene: C09G12.9 A; Map position: 4

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Db	240	SSSGGTIQADTIRASVMSAVEEKIRAKLRERMGTNSAEMASIRTTSDELREGQQKLKRML	299
Qу	275	TRLDQEVAEVDKNIELLKKKDEELSSALEKMENQSENNDIDEVIIPTAPLYKQILNLYAE : : : : : : : : ! : !:	334
Db	300	EELETQRSSLQTACEIYTAKKAELAKALSD-AGGTDAPPIDEAIDAAFPLHRQIVLNYAK	358
Qy	335	ENAIEDTIFYLGEALRRGVIDLDVFLKHVRLLSRKQFQLRALMQKARKTAGL 386 : :: :: : : : : :	
Db	359	DLTCQSLKKRQITLAEYLRHVRDVSREQFIYRATMQKCRRTAGL 402	

us-10-053-975a-1.rup

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on:

June 15, 2005, 10:20:07; Search time 168.534 Seconds

(without alignments)

1184.992 Million cell updates/sec

Title:

US-10-053-975A-1

Perfect score:

2047

Sequence:

1 MAVSESQLKKMVSKYKYRDL......FQLRALMQKARKTAGLSDLY 390

Scoring table:

BLOSUM62

Gapop 10.0, Gapext 0.5

Searched:

1612378 seqs, 512079187 residues

Total number of hits satisfying chosen parameters:

1612378

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 100 summaries

Database:

UniProt_03:*

1: uniprot_sprot:* 2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21	2047 2041 1945.5 1927.5 1908.5 1899 1705.5 1682.5 1024.5 976 754.5 749.5 599 522 435 435 429.5 427 415.5	100.0 99.7 95.0 94.2 93.8 83.3 82.2 50.0 47.7 36.6 29.3 25.2 21.3 21.0 20.9	390 390 391 391 391 392 390 394 395 408 402 425 326 249 114 398 379 580 249 174 476	121222222222222222222222222222222222222	T101_HUMAN Q9BUM5 T101_MOUSE Q6IRE4 Q7TSE5 Q918G8 Q6IQ70 Q6NUD5 Q6P2Z8 Q9VVA7 Q7Q6B6 O76258 Q6DDX9 Q8MQZ0 Q7T2M2 Q9LHG8 Q8IXO4 Q7S4R9 Q7TQD3 O08761 Q66KB7	Q99816 homo sapien Q9bum5 homo sapien Q61187 mus musculu Q6ire4 rattus norv Q7tse5 rattus norv Q9i8g8 chelonia my Q6iq70 brachydanio Q6nud5 xenopus lae Q6p2z8 xenopus tro Q9vva7 drosophila Q7q6b6 anopheles g Q6ddx9 xenopus lae Q8mqz0 drosophila Q7t2m2 ameiurus ne Q9lhg8 arabidopsis Q8ix04 homo sapien Q7s4r9 neurospora Q7tqd3 mus musculu Q66kb7 xenopus tro

					us-10-053-975a-1.rup		•
22	398	19.4	471	2	Q6DBY5	Q6d	lby5 brachydanio
23	396.5	19.4	402	2	Q6ESB7	Q6e	sb7 oryza sativ
24	391	19.1	368	2	Q9FFY6	q9f	fy6 arábidopsis
25	388	19.0	83	2	Q6LBE4	Q61	be4 mus musculu
26	371	18.1	177	2 2 2	Q96FF5	Q96	iff5 homo sapien
27	371	18.1	341	2	Q9NUX7		ux7 homo sapien
28	341	16.7	357	2 2 1	Q6P2F0	Q6p	2f0 homo sapien
29	303	14.8	385		ST22_YEAST		604 saccharomyc
30	301.5	14.7	378	2 2 2	Q6C148		:148 yarrowia li
31	301.5	14.7	378	2	Q873M7		'3m7 yarrowia li
32	294	14.4	111	2	Q8BU96		u96 mus musculu
33	294	14.4	468	2	Q6FS29		s29 candida gla
34	239	11.7	496	2	Q6BID5		id5 debaryomyce
35	215	10.5	445	2	Q75EU1		eu1 ashbya goss
36	169	8.3	376	2	P78998	P/8	8998 saccharomyc
37	155.5	7.6	299	2	Q6CS18	QGC	:s18 kluyveromyc
38 39	154.5 153.5	7.5 7.5	397 148	2	Q6NW27 Q6FIG7	Q011	w27 homo sapien ig7 homo sapien
40	153.3	7.5	283	2 2 . 2	Q641Z8		128 rattus norv
41	152	7.4	1158	2	Q6GM32		m32 xenopus lae
42	151.5	7.4	485	5	Q804G3		Ag3 brachydanio
43	148.5	7.3	148	2 2 2	Q6NXQ6		1xq6 homo sapien
44	148.5	7.3	148		09NZ81	09n	z81 homo sapien
45	148.5	7.3	397	2 2 2	Q8NEZ2	08n	nez2 homo sapien
46	148.5	7.3	397	2	Q96DL9		dl9 homo sapien
47	147.5	7.2	263	ī	Y737_DROME		3475 drosophila
48	147	7.2 7.2	397	2	Q8CHS8		hs8 mus musculu
49	145.5	7.1	1130	2	Q8IZL8	Q8i	zl8 homo sapien
50	144.5	7.1	743	2	Q9BU60		ou60 homo sapien
51	144.5	7.1	1048	2	Q96FT1		ift1 homo sapien
52	144.5	7.1	1130	2	Q6NTE6		ite6 homo sapien
53	144	7.0	483	2	Q7T <u>3</u> 91		391 brachydanio
54	144	7.0	488	1	ANX7_HUMAN		0073 homo sapien
55	143	7.0	664	2	Q6CDQ5	Q6C	dq5 yarrowia li
56	142	6.9	505	1	ANXB_HUMAN	P50	1995 homo sapien
57	142	6.9	1021	2	015451	012	451 homo sapien
58	141.5	6.9	284	2	Q6BGN1	QOD	ogn1 debaryomyce 7384 mus musculu
59	141.5	6.9 6.9	503	1.	ANXB_MOUSE		21f1 mus musculu
60 61	141.5 141	6.9	503 247	2	Q921F1 Q945K9		15k9 arabidopsis
62	141	6.9	463	2 1	ANX7_MOUSE	007	7076 mus musculu
63	141	6.9	463	Ž	Q922A2		22a2 mus musculu
64	140.5	6.9	1167	2	Q640S7		10s7 xenopus tro
65	139.5	6.8	900	2	Q6PSU8		su8 arabidopsis
66	139	6.8	415	2	Q8CCV9	080	cv9 mus musculu
67	139	6.8	1284	2	015450	o15	3450 homo sapien
68	138.5	6.8	827	2	060130)130 schizosacch
69	138.5	6.8	838	2	Q871B7		71b7 neurospor <u>a</u>
70	138.5	6.8	4903	1	MLL3_MOUSE		orh4 mus musculu
71	138	6.7	526	2	Q804G4)4g4 brachydanio
72	137	6.7	463	2	Q8BP75		p75 mus musculu
73	136.5	6.7	503	1	ANXB_RABIT		3477 oryctolagus
74	136	6.6	137	2	Q9CQ35	Qgo	cqj5 m mus muscu
75	136	6.6	1066	2	Q7XE90		ke90 oryza sativ
76	135.5	6.6	437	2	Q9STN0		stnO arabidopsis
77 -	135.5	6.6	895	2	Q9LIW7	Q9 I	liw7 oryza sativ
78 70	135.5	6.6	998 1105	2	Q6GNM2		nm2 xenopus lae mj4 mus musculu
79 80.	135.5 135	6.6 6.6	463	2	Q8BMJ4 Q8VIN2		/in2 rattus norv
80. 81	135	6.6	463	2	Q6IRJ7		irj7 rattus norv
82	135	6.6	674	2	Q18106		3106 caenorhabdi
83	135	6.6	703	2	Q7SCK8		sck8 neurospora
84	134.5	6.6	809	2	Q7ZVN7		zvn7 brachydanio
				_	• = · · · ·	• •	•

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us-10-053-975a-1.rup
                                                                               086316 mycobacteri
           134
                     6.5
                               198
                                          086316
                                      2 2
                               201
                                                                               Q7d8k1 mycobacteri
 86
           134
                                          Q7D8K1
                     6.5
                                                                               Q9utp6 schizosacch
Q641h0 xenopus lae
Q7u0b9 mycobacteri
                    6.5
                               485
502
 87
           134
                                          09UTP6
                                      2
 88
           134
                                          Q641H0
                    6.5
6.5
6.5
6.5
           133
 89
                               198
                                      2
                                          Q7U0B9
 90
                               548
                                          Q6DDW4
                                                                               Q6ddw4 xenopus lae
           133
                               471
512
        132.5
132.5
 91
                                          Q9LD31
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 92
                                                                               Q6nre3 xenopus lae
Q9z502 streptomyce
                                          Q6NRE3
 93
        132.5
                     6.5
                               671
                                          Q9Z502
 94
95
96
                                                                               Q7z429 homo sapien
Q94b77 arabidopsis
           132
                     6.4
                               371
                                          Q7Z429
        131.5
131.5
                     6.4
                               900
                                          Q94B77
                                                                               Q9fiu0 arabidopsis
Q8lcl8 arabidopsis
O59907 neurospora
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                     6.4
                                          Q9FIU0
 97
                     6.4
           131
                               101
                                          Q8LCL8
 98
                               472
                                          059907
           131
                     6.4
 99
           131
                     6.4
                               472
                                          Q6M9D4
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                     6.4
                               475
100
           131
                                          Q7RZ53
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RESULT 1
T101_HUMAN
                            STANDARD;
       T101_HUMAN
                                                 PRT:
                                                           390 AA.
ΙD
       Q99816;
AC
      16-OCT-2001 (Rel. 40, Created)
16-OCT-2001 (Rel. 40, Last sequence update)
25-OCT-2004 (Rel. 45, Last annotation update)
DT
DT
DT
       Tumor susceptibility gene 101 protein.
DE
      Name=TSG101:
GN
05
      Homo sapiens (Human).
      Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
0C
0C
0X
      NCBI_TaxID=9606:
       [1]
RN
RP
       SEQUENCE FROM N.A.
       TISSUE=Placenta;
RC
      MEDLINE=97148696; PubMed=9019400; DOI=10.1016/S0092-8674(00)81866-8;
RX
      Li L., Li X., Francke U., Cohen S.N.;
"The TSG101 tumor susceptibility gene is located in chromosome 11 band p15 and is mutated in human breast cancer.";
RA
RT
RT
       cell 88:143-154(1997).
RL
       [2]
RN
RP
       ERRATUM.
      MEDLINE=99080983; PubMed=9867424; DOI=10.1016/S0092-8674(00)89342-3;
RX
       Li L., Francke U., Cohen S.N.;
RA
       cell 93:660-660(1998).
RL
RN
RP
      ALTERNATIVE SPLICING (ISOFORM 2).
      MEDLINE=98031856; PubMed=9366528; DOI=10.1038/sj.onc.1201591;
RX
RA
       Gayther S.A., Barski P., Batley S.J., Li L., de Foy K.A., Cohen S.N.,
      Ponder B.A., Caldas C.;
"Aberrant splicing of the TSG101 and FHIT genes occurs frequently in multiple malignancies and in normal tissues and mimics alterations previously described in tumours.";
Oncogene 15:2119-2126(1997).
RA
RT
RT
RT
RL
RN
       [4]
RP
      ALTERNATIVE SPLICING.
      MEDLINE=97384943; PubMed=9242438;
RX
       Lee M.P., Feinberg A.P.;
"Aberrant splicing but not mutations of TSG101 in human breast
RA
RT
       cancer."
RT
      Cancer Res. 57:3131-3134(1997).
RL
```

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RN
RP
      ALTERNATIVE SPLICING.
RX
      MEDLINE=99054675; PubMed=9840940; DOI=10.1038/sj.onc.1202529;
      Wagner K.-U., Dierisseau P., Rucker E.B. III, Robinson G.W.,
RA
RA
      Hennighausen L.;
      "Genomic architecture and transcriptional activation of the mouse and
RT
      human tumor susceptibility gene TSG101: common types of shorter
RT
      transcripts are true alternative splice variants.'
RT
      Oncogene 17:2761-2770(1998).
RL
RN
      [6]
RP
      INTERACTION WITH DMAP1.
      MEDLINE=20347709; PubMed=10888872; DOI=10.1038/77023;
RX
      Rountree M.R., Bachman K.E., Baylin S.B.;
RA
      "DNMT1 binds HDAC2 and a new co-repressor, DMAP1, to form a complex at
RT
     replication foci.";
RT
      Nat. Genet. 25:269-277(2000).
RL
RN
      [7]
     STRUCTURE BY NMR OF 1-145.
MEDLINE=22001147; PubMed=12006492; DOI=10.1093/emboj/21.10.2397;
RP
RX
      Pornillos O., Alam S.L., Rich R.L., Myszka D.G., Davis D.R.,
RA
      Sundquist W.I.; "Structure and functional interactions of the Tsg101 UEV domain.";
RA
RT
      EMBO J. 21:2397-2406(2002).
RL.
RN
      STRUCTURE BY NMR OF 1-145.
RP
      MEDLINE=22289677; PubMed=12379843; DOI=10.1038/nsb856;
RX
      Pornillos O., Alam S.L., Davis D.R., Sundquist W.I.; "Structure of the Tsg101 UEV domain in complex with the PTAP motif of
RA
RT
      the HIV-1 p6 protein.
RT
      Nat. Struct. Biol. 9:812-817(2002).
RL
      -!- FUNCTION: May be involved in cell growth and differentiation. Acts
CC
CC
           as a negative growth regulator.
      -!- SUBUNIT: Interacts with ubiquitin, stathmin, GMCL and DMAP1 (By
CC
           similarity).
CC
      -!- SUBCELLULAR LOCATION: Mainly cytoplasmic. Depending on the stage of the cell cycle, detected in the nucleus.
CC
CC
      -!- ALTERNATIVE PRODUCTS:
CC
           Event=Alternative splicing; Named isoforms=2;
CC
             Comment=Additional isoforms seem to exist. Several shorter
CC
             isoforms are detected in primary breast cancers and other
CC
CC
             tumors:
CC
           Name=1;
             IsoId=Q99816-1; Sequence=Displayed;
CC
\mathsf{CC}
           Name=2;
             IsoId=Q99816-2; Sequence=VSP_004440;
CC
      Note=Detected in normal as well as cancer tissues;
-!- TISSUE SPECIFICITY: Heart, brain, placenta, lung, liver, skeletal,
cc
CC
      kidney and pancreas.
-!- DOMAIN: The UEV domain is required for the interaction of the
CC
CC
           complex with ubiquitin.
CC
      -!- DOMAIN: The coiled coil domain may interact with stathmin.
CC
CC
      -!- SIMILARITY: Contains 1 UEV (ubiquitin E2 variant) domain.
CC
      This SWISS-PROT entry is copyright. It is produced through a collaboration
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      between the Swiss Institute of Bioinformatics and the EMBL outstation - the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial
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      entities requires a license agreement (See http://www.isb-sib.ch/announce/
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      or send an email to license@isb-sib.ch).
CC
CC
      EMBL; U82130; AAC52083.1; -.
DR
DR
      PDB; 1KPP; NMR; A=1-145.
```

```
us-10-053-975a-1.rup
DR
     PDB; 1KPQ; NMR; A=1-145.
     PDB; 1M4P; NMR; A=1-145.
PDB; 1M4Q; NMR; A=1-145.
DR
DR
     IntAct; Q99816; -.
Genew; HGNC:15971; TSG101.
DR
DR
     H-InvDB; HIX0009490; -.
DR
     MIM; 601387:
DR
     GO: GO:0003677; F:DNA binding; TAS.
DR
     GO; GO:UUU307/; F:DNA DINGING; IAS.
GO; GO:0003714; F:transcription corepressor activity; TAS.
GO; GO:0006512; P:ubiquitin cycle; TAS.
InterPro; IPR010983; EF_Hand_like.
InterPro; IPR008883; Tsg101.
InterPro; IPR000608; UBQ_conjugat.
Pfam; PF05743; Tsg101; 1.
PROSITE; PS00183; UBIQUITIN_CONJUGAT_1; FALSE_NEG.
PROSITE; PS50127; UBIQUITIN_CONJUGAT_2; FALSE_NEG.
DR
DR
DR
DR
DR
DR
DR
DR
     3D-structure; Alternative splicing; Coiled coil; Growth regulation;
KW
     Protein transport; Transport; Ubl conjugation pathway.
KW
                          133
316
                                     UEV.
FT
     DOMAIN
                  235
                                     Coiled coil (Potential).
FT
     DOMAIN
                                     Missing (in isoform 2).
     VARSPLIC
                   15
                          119
FT
FΤ
                                     /FTId=VSP_004440.
                           11
FT
     HELIX
                           31
FT
     HELIX
                    33
                           34
     TURN
FT
                           44
47
FT
     STRAND
     TURN
FT
                           63
FT
     STRAND
                           65
75
FT
     TURN
                    64
FT
     STRAND
                    66
                   86
                           88
FT
     STRAND
     STRAND
                          103
                  103
FT
                  105
                          106
FT
     TURN
FT
     STRAND
                  108
                          109
                          115
FT
     HELIX
                  112
                  116
                          116
FT
     TURN
                  119
                          120
FT
     TURN
FT
     HELIX
                  124
                          137
     SEQUENCE
                 390 AA;
                           43944 MW;
                                     ADD6912FC22DF162 CRC64;
SQ ·
                            100.0%;
                                      Score 2047; DB 1; Pred. No. 2.1e-99;
                                                            Length 390;
  Query Match
                            100.0%; Pred. No. _
tive 0; Mismatches
  Best Local Similarity
  Matches 390; Conservative
                                                       0;
                                                            Indels
                                                                       0; Gaps
                                                                                    0:
             1 MAVSESQLKKMVSKYKYRDLTVRETVNVITLYKDLKPVLDSYVFNDGSSRELMNLTGTIP 60
Qy
               1 MAVSESQLKKMVSKYKYRDLTVRETVNVITLYKDLKPVLDSYVFNDGSSRELMNLTGTIP 60
Db
            61 VPYRGNTYNIPICLWLLDTYPYNPPICFVKPTSSMTIKTGKHVDANGKIYLPYLHEWKHP 120
Qy
               61 VPYRGNTYNIPICLWLLDTYPYNPPICFVKPTSSMTIKTGKHVDANGKIYLPYLHEWKHP 120
Db
           121 QSDLLGLIQVMIVVFGDEPPVFSRPISASYPPYQATGPPNTSYMPGMPGGISPYPSGYPP 180
Qy
               Db
           121 OSDLLGLIQVMIVVFGDEPPVFSRPISASYPPYQATGPPNTSYMPGMPGGISPYPSGYPP 180
           181 NPSGYPGCPYPPGGPYPATTSSQYPSQPPVTTVGPSRDGTISEDTIRASLISAVSDKLRW 240
Qу
               181 NPSGYPGCPYPPGGPYPATTSSQYPSQPPVTTVGPSRDGTISEDTIRASLISAVSDKLRW 240
Db
           241 RMKEEMDRAQAELNALKRTEEDLKKGHQKLEEMVTRLDQEVAEVDKNIELLKKKDEELSS 300
Qy
               241 RMKÉÉMDRÁQÁELNALKRTÉEDLKKGHQKLÉEMVTRLÓQEVÁEVDKNIELLKKKDEELSS 300
Db
```

us-10-053-975a-1.rup

Qy	301 ALEKMENQSENNDIDEVIIPTAPLYKQILNLYAEENAIEDTIFYLGEALRRGVIDLDVFL 360
Db	301 ALEKMENQSENNDIDEVIIPTAPLYKQILNLYAEENAIEDTIFYLGEALRRGVIDLDVFL 360
Qy	361 KHVRLLSRKQFQLRALMQKARKTAGLSDLY 390
Db	361 KHVRLLSRKQFQLRALMQKARKTAGLSDLY 390